=> d his

- (FILE 'HOME' ENTERED AT 10:06:37 ON 16 DEC 2008)
- FILE 'CA' ENTERED AT 10:06:47 ON 16 DEC 2008
- L1 1163 S POLYPHENYLENEETHYN? OR PHENYLENEETHYN? OR POLYPHENYLENE ETHYN? OR PHENYLENE (1A) ETHYN? OR POLYETHYN? PHENYLENE
- L2 494 S L1 AND (FLUORESC? OR LUMINESC? OR PHOTOLUMINESC?) OR ELECTROLUMINESC?)
- L3 10 S L1 AND CROWN?
- L4 76 S L2 AND (COMPLEX? OR CHELAT? OR PODAND OR CALIX?)
- L5 83 S L3-4
- L6 36 S L5 AND PY<2005
 - FILE 'BIOSIS' ENTERED AT 10:09:26 ON 16 DEC 2008
- L7 0 S L6
 - FILE 'MEDLINE' ENTERED AT 10:09:41 ON 16 DEC 2008
- L8 9 S L6
 - FILE 'CA, MEDLINE' ENTERED AT 10:10:07 ON 16 DEC 2008
- L9 37 DUP REM L6 L8 (8 DUPLICATES REMOVED)
- => d bib, ab 19 1-37
- L9 ANSWER 19 OF 37 CA COPYRIGHT 2008 ACS on STN
- AN 137:338473 CA
- TI Synthesis and properties of calix[4]arene-containing poly(phenyleneethynylene)s
- AU Wosnick, Jordan H.; Swager, Timothy M.
- CS Dept. of Chemistry, Massachusetts Inst. of Technology, Cambridge, MA, 02139, USA
- SO Polymer Preprints (American Chemical Society, Division of Polymer Chemistry) (2002), 43(2), 1001-1002
- AB The synthesis and properties of several calix[4]arene-substituted poly(phenyleneethynylene)s are reported. These polymers differ in their modes of attachment of the calix[4]arene group to the polymer backbone and in the mobility of the calix[4]arene skeleton. Polymers contg. a modified crown ether linkage connecting the calix[4]arene lower rim to the polymer main chain were extremely prone to gelation during polymn., a phenomenon not seen in polymers with calix[4]arenes attached via a single lower-rim tether or through an upper-rim linker group. The gelation effect is attributed to a zipper-like interdigitation of the rigidly held calix[4]arene groups on the growing polymer chains.
- L9 ANSWER 20 OF 37 CA COPYRIGHT 2008 ACS on STN
- AN 137:233185 CA
- TI Directing energy transfer within conjugated polymer films
- AU Rose, Aimee; Kim, Jinsang; McQuade, D. Tyler; Zhu, Zhennguo; Swager, Timothy M.
- CS Department of Chemistry, Massachusetts Institute of Technology, Cambridge, MA, 02139, USA
- SO PMSE Preprints (2002), 87, 377
- AB A striated multipolymer system was developed, which utilizes directional energy transfer to overcome the z-direction limitation in energy transfer within poly(p-phenylene ethynylene) conjugated polymers (CP). The polymers used are poly(p-phenylene ethynylene)s and copolymers of benzo-crown ethers and substituted benzodiacetylenes. The polymers have large spectral overlap between a donor emission and an acceptor absorption, ranging from the blue to the red, which encourages energy transfer from one polymer to a second, in LB multilayers. Two of the polymers were also designed to be non-aggregating and amphiphilic, thus allowing manipulation at the air-water interface. Energy can be preferentially transferred to the surface of a thin film by utilizing 8,16, 24, and 32 CP layers of sequentially decreasing band gap.

STN Columbus

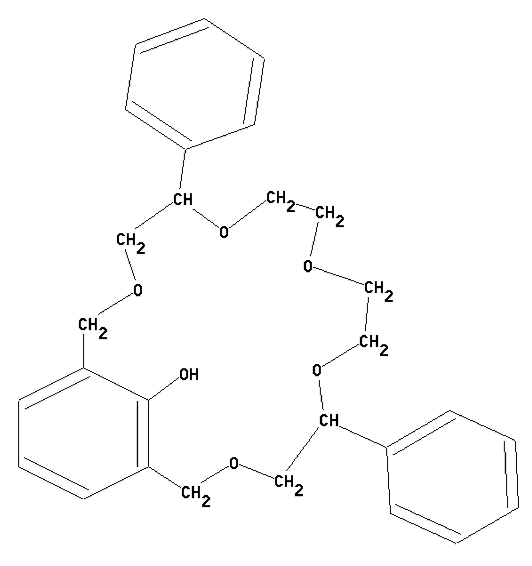
- AN 134:109803 CA
- TI Ion-specific aggregation in conjugated polymers: highly sensitive and selective fluorescent ion chemosensors
- AU Kim, Jinsang; McQuade, D. Tyler; McHugh, Sean K.; Swager, Timothy M.
- CS Dep. Chem., Massachusetts Inst. Technology, Cambridge, MA, 02139, USA
- SO Angewandte Chemie, International Edition (2000), 39(21), 3868-3872
- AB A new transduction mechanism based on the aggregation of conjugated sensory polymers induced by K+ ions is reported; this new system displays enhanced sensitivity because of energy migration processes and has a high selectivity for K+ over Na+ ions. The poly(p-phenylene ethynylene)s were synthesized by the Sonogashira-Hagihara coupling reaction.
- L9 ANSWER 31 OF 37 CA COPYRIGHT 2008 ACS on STN
- AN 132:317347 CA
- TI A poly(phenyleneethynylene) K+ chemosensor: detection via intermolecular aggregation
- AU Kim, Jinsang; McQuade, D. Tyler; McHugh, Sean K.; Swager, Timothy M.
- CS Department of Chemistry, Massachusetts Institute of Technology, Cambridge, MA, 02139, USA
- SO Polymer Preprints (American Chemical Society, Division of Polymer Chemistry) (2000), 41(1), 32-33
- AB Poly(phenyleneethynylene)s with 15-crown-5 side groups on every other repeating unit selectively detects K' by forming intermol. bridges. The extent of the aggregations can be tuned by controlling the steric bulk of side groups.
- L9 ANSWER 32 OF 37 CA COPYRIGHT 2008 ACS on STN
- AN 131:299764 CA
- TI High-Spin Polyphenoxyl Based on Poly(1,4-phenyleneethynylene)
- AU Nishide, Hiroyuki; Maeda, Tadatoshi; Oyaizu, Kenichi; Tsuchida, Eishun
- CS Department of Polymer Chemistry, Waseda University, Tokyo, 169-8555, Japan
- SO Journal of Organic Chemistry (1999), 64(19), 7129-7134
- AB Rodlike poly(1,4-phenyleneethynylene) 2-substituted with multiple pendant phenoxyls 1 was synthesized by polymg. 4-bromo-2-(3,5-di-tert-butyl-4-hydroxyphenyl)ethynylbenzene 10a using the catalyst of a palladium-triphenylphosphine complex and cuprous iodide and subsequent heterogeneous oxidn. The corresponding dimer 2 was also synthesized; X-ray anal. of its precursor 4 indicated a linear phenyleneethynylene backbone and twisted dihedral angles of 50
- and 77° for the pendant phenol groups. ESR spectra suggested a delocalized spin distribution from the pendant phenoxyl to the backbone. The diphenoxyl 2 had a triplet (S = 2/2) ground state. The spin concn. of the polyphenoxyl 1 could not be increased beyond 0.7 spin/unit due to its low solvent soly.; 1 with a spin concn. of 0.62 had an av. S of 3/2.
- L9 ANSWER 33 OF 37 CA COPYRIGHT 2008 ACS on STN
- AN 131:177205 CA
- TI Photophysics of metal-organic π -conjugated oligomers and polymers
- AU Ley, K. D.; Walters, K. A.; Schanze, K. S.
- ${\tt CS}$ Department of Chemistry, University of Florida, Gainesville, FL, 32611-7200, USA
- SO Synthetic Metals (1999), 102(1-3), 1585-1586
- AB A series of phenylene-ethynylene based π -conjugated oligomers that contain a 2,2'-bipyridine metal chelating unit has been synthesized by using Pd-mediated coupling chem. The photophysics of the free oligomers and complexes of the oligomers with the ReI(CO)3Cl metal chromophore is reported. These oligomers serve as excellent models for π -conjugated metal-org, polymers.
- => log y STN INTERNATIONAL LOGOFF AT 10:10:48 ON 16 DEC 2008

=> d his

(FILE 'HOME' ENTERED AT 16:44:28 ON 15 DEC 2008) FILE 'REGISTRY' ENTERED AT 16:44:49 ON 15 DEC 2008 L1STRUCTURE UPLOADED 4 S L1 L2 L3 118 S L1 FULL L48 S L3 AND (POLYMER OR ETHYN?) FILE 'CA' ENTERED AT 16:48:33 ON 15 DEC 2008 L5 16 S L3 66 S PSEUDO (1W)CROWN L6 L7 76 S L5-6

=>

Uploading C:\Program Files\Stnexp\Queries\crown.str



=> d 14 1-8

L4 ANSWER 1 OF 8 REGISTRY COPYRIGHT 2008 ACS on STN

RN 864920-14-9 REGISTRY

CN 3,6,9,12,15-Pentaoxabicyclo[15.3.1]heneicosa-1(21),17,19-trien-21-ol, 19-[[2-[[2,5-bis(decyloxy)-4-iodophenyl]ethynyl]-5-ethynylphenyl]ethynyl]-5,13-diphenyl-, (5S,13S)-, homopolymer (9CI) (CA INDEX NAME) MF (C66 H79 I O8)x

Absolute stereochemistry.

1 REFERENCES IN FILE CA (1907 TO DATE)

L4 ANSWER 2 OF 8 REGISTRY COPYRIGHT 2008 ACS on STN

RN 864920-13-8 REGISTRY

CN 3,6,9,12,15-Pentaoxabicyclo[15.3.1]heneicosa-1(21),17,19-trien-21-ol, 19-[2-[2-[2-[2,5-bis(decyloxy)-4-iodophenyl]ethynyl]-5-ethynylphenyl]ethynyl]-5,13-diphenyl-, (5S,13S)- (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 3,6,9,12,15-Pentaoxabicyclo[15.3.1]heneicosa-1(21),17,19-trien-21-ol, 19-[[2-[[2,5-bis(decyloxy)-4-iodophenyl]ethynyl]-5-ethynylphenyl]ethynyl]-5,13-diphenyl-, (5S,13S)- (9CI)

MF C66 H79 I O8

Absolute stereochemistry.

1 REFERENCES IN FILE CA (1907 TO DATE)

L4 ANSWER 3 OF 8 REGISTRY COPYRIGHT 2008 ACS on STN

RN 864920-12-7 REGISTRY

CN 3,6,9,12,15-Pentaoxabicyclo[15.3.1]heneicosa-1(21),17,19-trien-21-ol, 19-[2-[2-[2-[2,5-bis(decyloxy)-4-iodophenyl]ethynyl]-5-[2-[tris(1-

methylethyl)silyl]ethynyl]phenyl]=5,13-diphenyl-, (5S,13S)- (CA INDEX NAME)
OTHER CA INDEX NAMES:

CN 3,6,9,12,15-Pentaoxabicyclo[15.3.1]heneicosa-1(21),17,19-trien-21-ol, 19-[[2-[[2,5-bis(decyloxy)-4-iodophenyl]ethynyl]-5-[[tris(1-

methylethyl)silyl]ethynyl]phenyl]ethynyl]-5,13-diphenyl-, (5S,13S)- (9CI)

MF C75 H99 I O8 Si

Absolute stereochemistry.

1 REFERENCES IN FILE CA (1907 TO DATE)

L4 ANSWER 4 OF 8 REGISTRY COPYRIGHT 2008 ACS on STN

RN 864920-11-6 REGISTRY

CN 3,6,9,12,15-Pentaoxabicyclo[15.3.1]heneicosa-1(21),17,19-trien-21-o1, 19-[2-ethynyl-5-[2-[tris(1-methylethyl)silyl]ethynyl]phenyl]ethynyl]-5,13-diphenyl-, (5S,13S)- (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 3,6,9,12,15-Pentaoxabicyclo[15.3.1]heneicosa-1(21),17,19-trien-21-ol, 19-[[2-ethynyl-5-[[tris(1-methylethyl)silyl]ethynyl]phenyl]ethynyl]-5,13-diphenyl-, (5S,13S)-(9CI)

MF C49 H56 O6 Si

Absolute stereochemistry.

1 REFERENCES IN FILE CA (1907 TO DATE)

L4 ANSWER 5 OF 8 REGISTRY COPYRIGHT 2008 ACS on STN

RN 864920-10-5 REGISTRY

CN 3,6,9,12,15-Pentaoxabicyclo[15.3.1]heneicosa-1(21),17,19-trien-21-ol, 5,13-diphenyl-19-[2-[2-(trimethylsilyl)ethynyl]-5-[2-[tris(1-methylethyl)silyl]ethynyl]phenyl]ethynyl]-, (5S,13S)- (CA INDEX

NAME)

OTHER CA INDEX NAMES:

CN 3,6,9,12,15-Pentaoxabicyclo[15.3.1]heneicosa-1(21),17,19-trien-21-ol, 5,13-diphenyl-19-[[2-[(trimethylsilyl)ethynyl]-5-[[tris(1-methylethyl)silyl]ethynyl]phenyl]ethynyl]-, (5S,13S)- (9CI) MF C52 H64 O6 Si2

Absolute stereochemistry.

1 REFERENCES IN FILE CA (1907 TO DATE)

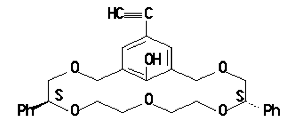
L4 ANSWER 6 OF 8 REGISTRY COPYRIGHT 2008 ACS on STN

RN 864920-05-8 REGISTRY

CN 3,6,9,12,15-Pentaoxabicyclo[15.3.1]heneicosa-1(21),17,19-trien-21-ol, 19-ethynyl-5,13-diphenyl-, (5S,13S)- (CA INDEX NAME)

MF C30 H32 O6

Absolute stereochemistry.



1 REFERENCES IN FILE CA (1907 TO DATE)

L4 ANSWER 7 OF 8 REGISTRY COPYRIGHT 2008 ACS on STN

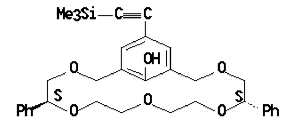
RN 864920-04-7 REGISTRY

CN 3,6,9,12,15-Pentaoxabicyclo[15.3.1]heneicosa-1(21),17,19-trien-21-ol, 5,13-diphenyl-19-[2-(trimethylsilyl)ethynyl]-, (5S,13S)- (CA INDEX NAME) OTHER CA INDEX NAMES:

CN 3,6,9,12,15-Pentaoxabicyclo[15.3.1]heneicosa-1(21),17,19-trien-21-ol, 5,13-diphenyl-19-[(trimethylsilyl)ethynyl]-, (5S,13S)- (9CI)

MF C33 H40 O6 Si

Absolute stereochemistry.



1 REFERENCES IN FILE CA (1907 TO DATE)

L4 ANSWER 8 OF 8 REGISTRY COPYRIGHT 2008 ACS on STN

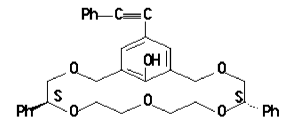
RN 666837-00-9 REGISTRY

CN 3,6,9,12,15-Pentaoxabicyclo[15.3.1]heneicosa-1(21),17,19-trien-21-ol, 5,13-diphenyl-19-(2-phenylethynyl)-, (5S,13S)- (CA INDEX NAME) OTHER CA INDEX NAMES:

CN 3,6,9,12,15-Pentaoxabicyclo[15.3.1]heneicosa-1(21),17,19-trien-21-ol, 5,13-diphenyl-19-(phenylethynyl)-, (5S,13S)- (9CI)

MF C36 H36 O6

Absolute stereochemistry. Rotation (+).



1 REFERENCES IN FILE CA (1907 TO DATE)

=> d bib, ab 17 1-76

L7 ANSWER 22 OF 76 CA COPYRIGHT 2008 ACS on STN

AN 143:338699 CA

TI Fluorescent molecular wire and its use in chiral sensor

IN Tobe, Yoshito; Hirose, Keiji

PA Japan Science and Technology Agency, Japan

SO Jpn. Kokai Tokkyo Koho, 29 pp.

CODEN: JKXXAF

DT Patent

LA Japanese

FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PΤ	JP 2005255778	7	20050922	JP 2004-67226	20040310
PI		A		JP 2004-67226	20040310
	JP 3950117	B2	20070725	TTO 0004 TD13635	00041100
	WO 2005087835	A1	20050922	WO 2004-JP17675	20041122
	US 20070179272	A1	20070802	US 2006-591920	20060907
PRAI	JP 2004-67226	A	20040310		

AB The wire has a fluorescent polymer main chain to which an optically active substituent from a 3,4,5-substituted Ph group of the structure 3-(R100CR4R5CR2R30)-5-(R110CR8R9CR6R6O)-4-R10C6H2 (R1 = H, C1-10 alkyl; R2-R9 = H, C1-30 linear alkyl, C2-30 branched alkyl, C3-30 cycloalkyl, C6-30 aryl, C7-30 aralkyl; R3 and R4, R7 and R8 may be linked to form C2-60 alkylene; R10, R11 = H, C1-15 alkyl optionally contg. heteroatom; and R10 and R11 may be linked to form C2-30 alkylene) is conjugatively linked. The wire has high sensitivity and improved asym. recognition and is suitable for primary amine chiral sensor.

L7 ANSWER 24 OF 76 CA COPYRIGHT 2008 ACS on STN

AN 143:221498 CA

TI Preparation and evaluation of a chiral stationary phase covalently bound with chiral pseudo-18-crown-6 ether having 1-phenyl-1,2-cyclohexanediol as a chiral unit AU Hirose, Keiji; Yongzhu, Jin; Nakamura, Takashi; Nishioka, Ryota; Ueshige, Tetsuro; Tobe, Yoshito

CS Division of Frontier Materials Science, Graduate School of Engineering Science, Osaka University, Toyonaka, Osaka, 560-8531, Japan

SO Journal of Chromatography, A (2005), 1078(1-2), 35-41

AB A chiral stationary phase (CSP) was prepd. by chem. bonding a chiral pseudo-18-crown-6 type host having a 1-phenyl-1,2-cyclohexanediol unit to 3-aminopropyl silica gel. The chiral column was prepd. by the slurry-packing method in a stainless steel HPLC column. Normal mobile phases can be used with this CSP in contrast to conventional dynamic coating type CSPs. Enantiomers of 20 out of 30 amino compds., including 20 amino acids, 2 amino acid Me esters, 6 amino alcs., and 2 lipophilic amines, were efficiently sepd. on columns with this CSP. It is noteworthy that 15 amino compds. out of 30 were sepd. with better sepn. factors and shorter retention times compared to the corresponding CSP having pseudo-18-crown-6 with 1-phenyl-1,2-ethanediol as a chiral unit. In view of the correlation between the enantiomer selectivities obsd. in chromatog. and those obtained in gas phase FABMS-EL methods and soln. phase titrns., chiral recognition in the host-guest interaction likely contributes to enantiomer sepn.

- L7 ANSWER 27 OF 76 CA COPYRIGHT 2008 ACS on STN
- AN 141:431425 CA
- TI Selective and Sensitive Fluorescent Sensors for Metal Ions Based on Manipulation of Side-Chain Compositions of Poly(p-phenyleneethynylene)s
- AU Chen, Zhen; Xue, Cuihua; Shi, Wei; Luo, Fen-Tair; Green, Sarah; Chen, Jian; Liu, Haiying
- CS Department of Chemistry, Michigan Technological University, Houghton, MI, 49931, USA
- SO Analytical Chemistry (2004), 76(21), 6513-6518
- AB The syntheses and metal-responsive properties of poly(p-phenyleenethynylene)s with grafted new pseudo-crown-ether groups are reported. These polymers exhibit high sensitivities to alkali ions because of their collective optical properties, which are very sensitive to ion-induced conformational changes. The quenching of polymer fluorescence caused by the conformational changes is proportional to the ion concn. The selectivity of the sensing materials toward Li+ ions is significantly enhanced by controlling the size of the binding site via manipulation of the polymer side-chain compns. The polymers are very stable for their six-month solid-state storage at room temp.
- L7 ANSWER 30 OF 76 CA COPYRIGHT 2008 ACS on STN
- AN 140:235760 CA
- TI Chiral sensor
- IN Tobe, Yoshito; Hirose, Keiji
- PA Japan Science and Technology Corporation, Japan
- SO PCT Int. Appl., 29 pp.

ΡI	WO	2004018447	A1	20040304	WO 2003-JP7313	20030610
	US	20050227366	A1	20051013	US 2005-525012	20050217
	US	7358403	B2	20080415		
PRAI	JΡ	2002-239777	A	20020820		

AB This document discloses an optically active compd. having an unsatd. bond at the optically active combining site, wherein the unsatd. bond and a fluorescent substituent or a substituent capable of imparting fluorescence are united in a conjugated state; and a chiral sensor consisting of the compd. The chiral sensor can recognize specific chiral compds. highly sensitively and highly selectively.

- L7 ANSWER 34 OF 76 CA COPYRIGHT 2008 ACS on STN
- AN 138:313549 CA
- TI Preparation and evaluation of novel chiral stationary phases covalently bound with chiral pseudo-18-crown-6 ethers
- AU Hirose, Keiji; Nakamura, Takashi; Nishioka, Ryota; Ueshige, Tetsuro; Tobe, Yoshito
- CS Faculty of Engineering Science, Department of Chemistry, Osaka University, Toyonaka, Osaka, 560-8531, Japan
- SO Tetrahedron Letters (2003), 44(8), 1549-1551
- AB Novel chiral stationary phases consisting of silica gel covalently bound with chiral pseudo-18-crown-6 type hosts, which possess either an OH or OMe group as a binding functionality, were prepd. for enantiomer-sepn. of lipophilic amines.

- AN 138:169671 CA
- TI Chiral recognition of secondary amines by using chiral crown ether and podand AU Hirose, Keiji; Fujiwara, Akihito; Matsunaga, Kazuhisa; Aoki, Nobuaki; Tobe, Yoshito
- CS Faculty of Engineering Science, Department of Chemistry, Osaka University, and CREST, Japan Science and Technology Corporation (JST), Toyonaka, Osaka, 560-8531, Japan
- SO Tetrahedron Letters (2002), 43(47), 8539-8542
- AB Chiral crown ether (S,S)-3 having a pseudo-24-crown-8 ring and chiral podand (R,R)-4 were prepd. and both exhibited good chiral recognition ability toward secondary amines, N-.alpha.-dimethylbenzylamine (15) and propranolol (16).
- L7 ANSWER 41 OF 76 CA COPYRIGHT 2008 ACS on STN
- AN 136:14888 CA
- TI Stationary phase for chromatography
- IN Nishioka, Ryota; Ueshige, Tetsuro; Tobe, Yoshihito; Hirose, Keiji
- PA Sumika Chemical Analysis Service Ltd., Japan
- SO Jpn. Kokai Tokkyo Koho, 6 pp.
- PI JP 2001327863 A 20011127 JP 2000-153051 20000524
- PRAI JP 2000-153051 20000524
- AB The title stationary phase is made by bonding a crown ether of specific structure to silica gel carrier. The phase has a structure as shown in graph (I), where R1, R2, and R3 are monovalent groups with at least one of them bonded to Si atom. R4 is H atom or an alkyl group of 1-10 C. R5-12 are H atom or substituted alkyl groups of 1-30 C or cyclic alkyl, aryl groups. The better results are given while R6-R7 or R10-R11 are bonded each other to form alkylene groups of 2-60 C. N is an integer of 1-10 and X is -NHCO- or NH- group. The stationary phase has good durability and sepn. property.
- L7 ANSWER 45 OF 76 CA COPYRIGHT 2008 ACS on STN
- AN 135:231767 CA
- ${\tt TI}$ Preparation and evaluation of novel chiral stationary phases chemically bonded with chiral pseudo crown ether
- AU Nishioak, R.; Ueshige, T.; Nakamura, T.; Hirose, K.; Tobe, Y.
- CS Sumika Chemical Analysis Service, Japan
- SO Chromatography (2000), 21(4), 294-295
- LA Japanese
- AB We have developed novel chiral stationary phases (CSPs) covalently bonded with chiral pseudo crown ether contg. Ph groups as chiral barrier which has high ability of discriminating enantiomers. These CSPs are chem. stable, so that both the reversed and the normal phases can be used. These are very effective for enantiomer sepns. of wide range of chiral amines, aminoalcs. and amino acids, esp. for hydrophobic amines.
- L7 ANSWER 46 OF 76 CA COPYRIGHT 2008 ACS on STN
- AN 135:191397 CA
- TI Enantiomeric separations of stimulant materials using chiral stationary phase bonded with pseudo crown ether
- AU Ueshige, T.; Nishioka, R.; Nakamura, T.; Hirose, K.; Tobe, Y.
- CS Sumika Chemical Analysis Service, Japan
- SO Chromatography (2000), 21(4), 368-369
- LA Japanese
- AB We have developed novel chiral stationary phases (CSPs) chem. bonded with chiral pseudo 18-crown-6 ether. These CSPs are effective for enantiomer sepns. of chiral amines, amino acids, and esp. of amino alcs. In this report, the method for direct chiral sepns. of DL-norephedrine, one of stimulant materials, and its related compds. with HPLC using these CSPs were developed. This method is suitable for practical use.

- AN 133:349944 CA
- TI Enantioselective complexation of phenolic crown ethers with chiral aminoethanol derivatives: effects of substituents of aromatic rings ofhosts and guests on complexation
- AU Hirose, Keiji; Ogasahara, Kazuko; Nishioka, Kazuyuki; Tobe, Yoshito; Naemura, Koichiro
- CS Faculty of Engineering Science, Department of Chemistry, Osaka University, Toyonaka, Osaka, 560-8531, Japan
- SO Perkin 2 (2000), (9), 1984-1993
- AB Optically active azophenolic crown ethers having Ph groups substituted at the resp. para-position were prepd. and their assocn. consts. with chiral aminoethanol derivs., including 2-amino-2-phenylethanols having an electron-donating or an electron-withdrawing group, were detd. in chloroform by means of UV-vis titrn. methods. The enantioselectivities of these crown ethers are estd. from the ratio of the assocn. consts. KR/KS and the effect of arom. substituents of both hosts and guests on the binding abilities and enantioselectivities is discussed. The structures of the complexes were investigated on the basis of the 1H NMR and UV-vis spectra.
- L7 ANSWER 56 OF 76 CA COPYRIGHT 2008 ACS on STN
- AN 129:95231 CA
- OREF 129:19643a,19646a
- TI Preparation and temperature-dependent enantioselectivities of homochiral phenolic crown ethers having aryl chiral barriers: thermodynamic parameters for enantioselective complexation with chiral amines
- AU Naemura, Koichiro; Nishioka, Kazuyuki; Ogasahara, Kazuko; Nishikawa, Yasushi; Hirose, Keiji; Tobe, Yoshito
- CS Dep. Chem., Fac. Eng. Sci., Osaka Univ., Osaka, 560, Japan
- SO Tetrahedron: Asymmetry (1998), 9(4), 563-574
- AB (In this abstr. 1-4 = I-IV, resp.). Homochiral crown ether (S,S)-1 contg. 1-naphthyl groups as chiral barriers together with the phenol moiety was prepd. by using (S)-3 as a chiral subunit which was resolved in enantiomerically pure form by lipase-catalyzed enantioselective acylation of (.+-.)-3. Homochiral phenolic crown ether (S,S)-2, contg. Ph groups as chiral barriers, was also prepd. from (S)-4 which was derived from (S)-mandelic acid. The assocn. consts. for their complexes with chiral amines in CHCl3 were detd. at various temps. by the UV-visible spectroscopic method demonstrating that the crown ethers (S,S)-1 and (S,S)-2 displayed the large Δ R-S Δ G values of 6.2 and 6.4 kJ mol-1, resp., towards the amine (R)-2-amino-2-phenylethanol at 15.degree.C. Thermodn. parameters for complex formation were also detd. and a linear correlation between T Δ R-S Δ S and Δ R-S Δ H values was obsd.
- L7 ANSWER 58 OF 76 CA COPYRIGHT 2008 ACS on STN
- AN 126:171153 CA
- OREF 126:33069a,33072a
- TI Temperature dependent reversal of enantiomer selectivity in the complexation of optically active phenolic crown ethers with chiral amines
- AU Naemura, Koichiro; Fuji, Junichi; Ogasahara, Kazuko; Hirose, Keiji; Tobe, Yoshito
- CS Dep. Chem., Fac. Eng. Sci., Osaka Univ., Osaka, 560, Japan
- SO Chemical Communications (Cambridge) (1996), (24), 2749-2750
- AB Phenolic crown ethers (S,S)-I (R= 1-adamantyl, Ph, Me) and (R,R)-II were prepd. in enantiomerically pure forms; the enantiomer selectivities of crown ethers (S,S)-I (R= 1-adamantyl) and (R,R)-II in complexation with 2-aminopropan-1-ol reversed at ca. 6.degree. and increased with increasing temp. above the isoenantioselective temp.
- => log y